

**Claims after this response:**

1. (Previously Presented) A computer-implemented method for overlaying gene- or protein-related data on chromosome maps to provide at least one data-enhanced chromosome map as an output, said method comprising:
  - receiving said chromosome maps as a first input to a computer;
  - receiving a list comprising a plurality of gene- or protein-related data items as a second input to said computer, each data item including data other than data specifying a genetic location on said chromosome maps;
  - providing an identifier specifying a genetic location for each of said data items on said chromosome maps;
  - matching said identifiers with predefined identifiers on at least one of said chromosome maps;
  - reordering said data items to an order matching the order of said predefined identifiers on said at least one of said chromosome maps; and
  - displaying said data items on said at least one chromosome map at locations determined by said reordered data items to provide said data-enhanced output, wherein said receiving, matching, reordering and displaying are carried out automatically by said computer.
2. (Original) The method of claim 1, further comprising interactive selection by a user of at least one data type to be displayed during said displaying.
3. (Original) The method of claim 1, further comprising spatially grouping said gene- or protein-related data to correspond to spatial groupings of said associated genes on said at least one chromosome map.
4. (Original) The method of claim 1, further comprising compressing said gene- or protein-related data when required to display said gene- or protein-related data in an area in which all of the gene- or protein-related data cannot be discretely displayed.
5. (Original) The method of claim 1, further comprising zooming at least one of said gene- or protein-related data and said at least one chromosome map to display an enlarged view of additional detail relevant to a zoomed area.
6. (Original) The method of claim 1, further comprising querying and cutting information on the display that a user is not interested in viewing.

7. (Previously Presented) The method of claim 1, wherein said at least one chromosome map comprises a plurality of chromosome maps, said method further comprising maintaining focus and context of at least a portion of the display of said chromosome maps and gene- or protein-related data by displaying at least one chromosome map with overlaid gene- or protein-related data at a first magnification and at least one other chromosome map with overlaid gene- or protein-related data at a second magnification, so that said one map and said one other map are viewed simultaneously, said first magnification being different from said second magnification.

8. (Previously Presented) The method of claim 1, wherein said at least one chromosome map comprises a plurality of chromosome maps, said method further comprising maintaining focus and context of at least a portion of the display of said chromosome maps and gene- or protein-related data, by simultaneously displaying a high level view of all of said chromosome maps and gene- or protein-related data, a mid-level view displaying a magnified view of a selected portion of said high level view, and a detailed view displaying expanded, detailed information characterizing a selected portion of said mid-level view.

9. (Original) The method of claim 8, wherein said high-level view, mid-level view and detailed view are all interlinked so that changing one view automatically changes the other two views in the same way, substantially simultaneously.

10. (Original) The method of claim 1, further comprising displaying tooltips to display additional details relative to a selected portion of the display.

11. (Original) The method of claim 1, further comprising displaying popup dialogs to display additional details relative to a selected portion of the display.

12. (Previously Presented) The method of claim 1, further comprising accessing an external source of information relative to the data displayed, matching at least one of said identifiers with specific information in said external source; and displaying said specific information relative to said gene- or protein-related data associated with said at least one identifier, wherein said specific information comprises data other than markers, identifiers, or a chromosome map.

13. (Previously Presented) The method of claim 1, wherein said identifiers of said gene- or protein-related data are selected from published gene identifiers and symbols.

14. (Original) The method of claim 13, wherein said published gene identifiers and

symbols are selected from at least one of GenBank accession numbers, RefSeq accession numbers, UniGene Cluster ID's, UniGene ID's, official standard gene names, LocusLink ID, SwissProt ID's, and Protein Information Resource (PIR) ID's.

15. (Original) The method of claim 1, wherein said matching comprises providing a relational database which stores a set of cross-referenced tables for matching said identifiers with said predefined identifiers, and as the identifiers are read, they are matched with said predefined identifiers in the cross-referenced tables through standard database queries.

16. (Previously Presented) The method of claim 1, wherein said gene- or protein-related data comprises an expression matrix having rows and columns, wherein each said row of said matrix contains data values for a particular gene or protein across a set of measured samples, and results of each said measured sample are provided by data in respective columns of said matrix.

17. (Original) The method of claim 16, wherein said arbitrary gene- or protein-related data comprises a plurality of expression matrices.

18. (Previously Presented) The method of claim 1, wherein said arbitrary gene- or protein-related data comprises a matrix of at least one microarray of gene expression data, wherein each row of the matrix is associated with a particular gene, with data in the respective row being associated with said particular gene, and wherein said matching comprises reordering and spatial grouping of the rows based on matching the identifiers to the predefined identifiers.

19. (Original) The method of claim 18, wherein a visualization of the matrix resultant from said displaying comprises a heat map.

20. (Previously Presented) The method of claim 1, further comprising statistically assessing co-location values and displaying assessed co-location statistical significance along side said gene- or protein-related data.

21. (Previously Presented) The method of claim 1, further comprising the steps of:  
selecting additional information characterizing said gene- or protein-related data; and  
displaying said additional information along side of said display of the gene- or protein-related data and positioned relative to the respective locations on the chromosome map of the respective genes characterized by said gene- or protein-related data.

22. (Original) The method of claim 21, wherein said additional information comprises annotations.

23. (Original) The method of claim 22, wherein said annotations comprise gene ontology annotations.

24. (Original) The method of claim 21, wherein said additional information is selected from the group consisting of CGH data, protein levels, relevance scores and relevance densities.

25. (Previously Presented) The method of claim 22, wherein said gene- or protein-related data is displayed in matrix format and said additional information is displayed in at least one additional matrix.

26. (Previously Presented) The method of claim 21, wherein said gene- or protein-related data is displayed in scatter plot format.

27. (Previously Presented) The method of claim 1, wherein said gene- or protein- related data is imported from a plurality of experiments.

28. (Previously Presented) The method of claim 27, wherein said gene- or protein- related data is displayed with regard to each of the plurality of experiments on a single display.

29. (Original) The method of claim 21, wherein said additional information includes at least one of annotations, cellular localization of the genetic material, cluster data, and statistical data.

30. (Previously Presented) The method of claim 18, further comprising calculating row vectors of the values in the rows of the matrix; using an auxiliary process to obtain cluster data for said row vectors; and displaying said cluster data along side said display of said gene- or protein-related data.

31. (Previously Presented) The method of claim 30, wherein said matrix comprises a heat map, and wherein said cluster data and said gene- or protein-related data are displayed with color coding.

32. (Original) The method of claim 30, wherein said cluster data is displayed in a single column adjacent each matrix of gene- or protein-related data.

33. (Previously Presented) The method of claim 30, wherein said cluster data is displayed in a multi-column matrix adjacent each matrix of gene- or protein-related data.

34. (Previously Presented) The method of claim 1, wherein said gene- or protein-related data comprises a matrix of at least one microarray of gene expression data, wherein each row of the matrix is associated with a particular gene, and wherein each column of the matrix is

associated with a microarray experiment, wherein a portion of the total number of columns are associated with experiments taken from normal, healthy tissue, and another portion of the total number of columns are associated with experiments taken from tissue exhibiting an abnormality, said method further comprising dividing the matrix into two smaller matrices with a first matrix containing the columns associated with normal experiments and a second matrix containing the columns associated with abnormal experiments, and wherein said matching and displaying are performed with regard to both first and second matrices.

35. (Original) The method of claim 34, wherein the first and second matrices are displayed in color coding as heat maps.

36. (Original) The method of claim 34, further comprising calculating a relevance score for at least one row of the matrices by comparing expression values in the first matrix with expression values in the second matrix, and displaying at least one calculated relevance score along side the row to which each pertains.

37. (Original) The method of claim 36 wherein said calculating is interactively initiated via a user interface.

38. (Previously Presented) The method of claim 36, wherein the relevance score comprises a "p value" and the relevance score is displayed as a value calculated by  $(-\log p \text{ value})$ .

39. (Previously Presented) The method of claim 36, wherein a plurality of relevance scores is calculated and displayed as a line map.

40. (Previously Presented) The method of claim 36, wherein a plurality of relevance scores is calculated and displayed in color-coding as a heat map.

41. (Original) The method of claim 36, wherein relevance scores are calculated and displayed in a binary code.

42. (Previously Presented) The method of claim 36, wherein a plurality of relevance scores is calculated, said method further comprising defining a relevance density score based upon distances between genetic locations and relevance scores, and identifying chromosomal locations containing relevance density scores greater than or equal to the defined relevance density score.

43. (Previously Presented) The method of claim 36, wherein a plurality of relevance scores is calculated, said method further comprising filtering the relevance scores by setting at

least one relevance score limit value and displaying only those relevance scores which are greater than or equal to at least one relevance score limit value.

44. (Original) The method of claim 34, further comprising matching chromosomal copy number abnormality data with the gene-related data identifiers, and displaying the chromosomal copy number abnormality data along side the gene-related data to which each is matched.

45. (Original) The method of claim 44, wherein the chromosomal copy number abnormality data is displayed in third and fourth matrices, wherein each value in the third matrix is matched with the expression value in the first matrix having the same row and column location, and wherein each value in the fourth matrix is matched with the expression value in the second matrix having the same row and column location.

46. (Original) The method of claim 44, wherein the chromosomal copy number abnormality data is provided in columns interlaced with the columns of expression data in the first and second matrices.

47. (Original) The method of claim 44, wherein the chromosomal copy number abnormality is displayed in color-coding, as one or more heat maps.

48. (Original) The method of claim 44, wherein the chromosomal copy number abnormality data is displayed as one or more line maps.

49. (Original) The method of claim 45, further comprising calculating a relevance score for at least one row of the chromosomal copy number abnormality data by comparing chromosomal copy number abnormality values in the third matrix with chromosomal copy number abnormality values in the fourth matrix, and displaying at least one calculated relevance score along side the row to which each pertains.

50. (Previously Presented) The method of claim 49, wherein the relevance score comprises a "p value" and the relevance score is displayed as a value calculated by  $(-\log p \text{ value})$ .

51. (Previously Presented) The method of claim 49, wherein a plurality of relevance scores is calculated and displayed as a line map.

52. (Previously Presented) The method of claim 49, wherein a plurality of relevance scores is calculated and displayed in color-coding as a heat map.

53. (Previously Presented) The method of claim 49, wherein a plurality of relevance scores is calculated, said method further comprising defining a relevance density score based

upon distances between genetic locations and relevance scores, and identifying chromosomal locations containing relevance density scores greater than or equal to the defined relevance density score.

54. (Previously Presented) The method of claim 49, wherein a plurality of relevance scores is calculated, said method further comprising filtering the relevance scores by setting at least one relevance score limit value and displaying only those relevance scores which meet or exceed at least one relevance score limit value.

55. (Previously Presented) The method of claim 1, further comprising the steps of:  
selecting additional information related to one or more genes characterized by said gene- or protein-related data; and

displaying said additional information along side of said display of the gene- or protein-related data and positioned relative to the respective locations on the chromosome map of the respective genes characterized by said gene- or protein-related data.

56. (Original) The method of claim 55, wherein said additional information comprise at least one of polymorphism measurements, annotations, transcription factor binding sites, RNA expression values, allele information, alternative exon splicing data, mapping of CGH gene amplification/deletions, and protein abundance.

Claims 57-79 (Canceled).

80. (Previously Presented) A computer-implemented method for overlaying gene- or protein-related data on chromosome maps to provide a data-enhanced map, said method comprising:

- receiving said chromosome maps as a first input to a computer;
- receiving a list comprising a plurality of gene- or protein-related data items as a second input to said computer, each data item including data other than data specifying a genetic location on said chromosome maps;
- providing an identifier specifying a genetic location for each of said data items on said chromosome maps;
- matching said identifiers with predefined identifiers on at least one of said chromosome maps;
- reordering said data items to an order matching the order of said predefined identifiers on

said at least one of said chromosome maps; and

displaying said data items on said at least one chromosome map at locations determined by said reordered data items to provide said data enhanced map, wherein said receiving, matching and displaying are carried out automatically by said computer and wherein said gene- or protein-related data comprises a matrix of at least one microarray of gene expression data, wherein each row of said matrix is associated with a particular gene, and wherein each column of said matrix is associated with a microarray experiment, wherein a portion of the total number of columns is associated with experiments taken from normal, healthy tissue, and another portion of the total number of columns is associated with experiments taken from tissue exhibiting an abnormality,

said method further comprising dividing said matrix into two smaller matrices with a first matrix containing the columns associated with normal experiments and a second matrix containing the columns associated with abnormal experiments, and wherein said matching and displaying are performed with regard to both first and second matrices.

81. (Previously Presented) The method of claim 80, wherein the first and second matrices are displayed in color coding as heat maps.

82. (Previously Presented) The method of claim 80, further comprising calculating a relevance score for at least one row of the matrices by comparing expression values in the first matrix with expression values in the second matrix, and displaying at least one calculated relevance score along side the row to which each pertains.

83. (Previously Presented) The method of claim 82, wherein said calculating is interactively initiated via a user interface.

84. (Previously Presented) The method of claim 82, wherein the relevance score comprises a "p value" and the relevance score is displayed as a value calculated by  $(-\log p \text{ value})$ .

85. (Previously Presented) The method of claim 82, wherein a plurality of relevance scores is calculated and displayed as a line map.

86. (Previously Presented) The method of claim 82, wherein a plurality of relevance scores is calculated and displayed in color-coding as a heat map.

87. (Previously Presented) The method of claim 82, wherein relevance scores are calculated and displayed in a binary code.



88. (Previously Presented) The method of claim 82, wherein a plurality of relevance scores is calculated, said method further comprising defining a relevance density score based upon distances between genetic locations of genes to which said gene-related data are associated, and relevance scores of the gene-related data, and identifying chromosomal locations containing relevance density scores greater than or equal to the defined relevance density score.

89. (Previously Presented) The method of claim 82, wherein a plurality of relevance scores are calculated, said method further comprising filtering the relevance scores by setting at least one relevance score limit value and displaying only those relevance scores which are greater than or equal to at least one relevance score limit value.

90. (Previously Presented) The method of claim 80, further comprising matching chromosomal copy number abnormality data with the gene-related data identifiers, and displaying the chromosomal copy number abnormality data along side the gene-related data to which each is matched.

91. (Canceled)

92. (Previously Presented) The method of claim 90, wherein the chromosomal copy number abnormality data is provided in columns interlaced with the columns of expression data in the first and second matrices.

93. (Previously Presented) The method of claim 90, wherein the chromosomal copy number abnormality is displayed in color-coding, as one or more heat maps.

94. (Previously Presented) The method of claim 90, wherein the chromosomal copy number abnormality data is displayed as one or more line maps.

95. (Previously Presented) The method of claim 101, further comprising calculating a relevance score for at least one row of the chromosomal copy number abnormality data by comparing chromosomal copy number abnormality values in the third matrix with chromosomal copy number abnormality values in the fourth matrix, and displaying at least one calculated relevance score along side the row to which each pertains.

96. (Previously Presented) The method of claim 95, wherein the relevance score comprises a "p value" and the relevance score is displayed as a value calculated by  $(-\log p \text{ value})$ .

97. (Previously Presented) The method of claim 95, wherein a plurality of relevance scores is calculated and displayed as a line map.

98. (Previously Presented) The method of claim 95, wherein a plurality of relevance scores is calculated and displayed in color-coding as a heat map.

99. (Previously Presented) The method of claim 95, wherein a plurality of relevance scores is calculated, said method further comprising defining a relevance density score based upon distances between genetic locations and relevance scores, and identifying chromosomal locations containing relevance density scores greater than or equal to the defined relevance density score.

100. (Previously Presented) The method of claim 95, wherein a plurality of relevance scores is calculated, said method further comprising filtering the relevance scores by setting at least one relevance score limit value and displaying only those relevance scores which meet or exceed at least one relevance score limit value.

101. (Previously Presented) A computer facilitated method for overlaying gene- or protein-related data on chromosome maps, said method comprising:

receiving a list comprising a plurality of gene- or protein-related data items as an input to a computer, each data item having identifiers specifying a genetic location for each of said data items on said chromosome maps;

matching the identifiers with predefined identifiers on at least one of the chromosome maps;

displaying the data items on the at least one chromosome map where the genes associated with the respective data items are located, wherein said receiving, matching and displaying are carried out automatically by said computer, and wherein said gene- or protein-related data comprises a matrix of at least one microarray of gene expression data, wherein each row of the matrix is associated with a particular gene, and wherein each column of the matrix is associated with a microarray experiment, wherein a portion of the total number of columns are associated with experiments taken from normal, healthy tissue, and another portion of the total number of columns are associated with experiments taken from tissue exhibiting an abnormality,

dividing the matrix into two smaller matrices with a first matrix containing the columns associated with normal experiments and a second matrix containing the columns associated with abnormal experiments, wherein said matching and displaying are performed with regard to both first and second matrices; and

matching chromosomal copy number abnormality data with the gene-related data identifiers, and displaying the chromosomal copy number abnormality data along side the gene-related data to which each is matched, wherein the chromosomal copy number abnormality data is displayed in third and fourth matrices, wherein each value in the third matrix is matched with the expression value in the first matrix having the same row and column location, and wherein each value in the fourth matrix is matched with the expression value in the second matrix having the same row and column location.